First Hit

Generate Collection Print

L5: Entry 1 of 9

File: PGPB

Mar 25, 2004

DOCUMENT-IDENTIFIER: US 20040058417 A1

TITLE: Covalent joining of DNA to RNA by vaccinia topoisomerase and uses thereof

Detail Description Paragraph:

[0059] A DNA tag sequence can be attached to the isolated full-length mRNA using the methods described above. A preferred DNA tag sequence is shown in FIG. 11 both as a double stranded DNA cleavage substrate and as a covalent topoisomerase-DNA intermediate. The complementary strand of the topoisomerase-DNA intermediate includes a 3' overhang of from 1 to 4 nucleotides, which can be any mixture of adenine, guanine, cytosine or thymine, designated in the figure as N. These nucleotides will base pair with the first 1 to 4 bases of the 5' end of the isolated mRNA molecule, allowing the covalently attached topoisomerase to catalyze the transesterification reaction which joins the DNA tag to the end of the RNA sequence. The DNA tag sequence comprises a topoisomerase recognition site, preferably CCCTT, and in addition may comprise a recognition site for a site-specific restriction endonuclease, such as EcoR1, useful for the subsequent insertion of a cDNA molecule into an expression vector.

CLAIMS:

58. A method of claim 45, wherein the DNA $\underline{\text{tag}}$ sequence comprises a $\underline{\text{recognition site}}$ for a type I $\underline{\text{topoisomerase}}$.

tags or epitope tags. Affinity purification tags are generally peptide sequences that can interact with a binding partner immobilized on a solid support. Synthetic DNA sequences encoding multiple consecutive single amino acids, such as histidine, when fused to the expressed protein, may be used for one-step purification of the recombinant protein by high affinity binding to a resin column, such as nickel sepharose. An endopeptidase recognition sequence can be engineered between the polyamino acid tag and the protein of interest to allow subsequent removal of the leader peptide by digestion with Enterokinase, and other proteases. Sequences encoding peptides such as the chitin binding domain (which binds to chitin), glutathione-S-transferase (which binds to glutathione), biotin (which binds to avidin and strepavidin), and the like can also be used for facilitating purification of the protein of interest. The affinity purification tag can be separated from the protein of interest by methods well known in the art, including the use of inteins (protein self-splicing elements, Chong, et al, Gene 192:271-281, 1997).

Detailed Description Text (64):

The strand transfer reaction pathway is diagrammed in FIG. 10A. The biotinylated DNA Substrate which contains a single topoisomerase recognition site is immobilized on the Dynabeads (Dynal) streptavidin solid support. The biotin moiety (indicated by the black square) is introduced at the 5' end of the CCCTT-containing strand via standard protocols for automated oligonucleotide synthesis. The purified vaccinia topoisomerase is reacted with the bead-bound DNA to form a covalent enzyme-DNA donor complex, as illustrated. Enzyme not bound to DNA is removed by washing the beads with buffer. The strand transfer reaction is initiated by addition of the [.sup.32 P]-CMP labeled T7 transcript which is dephosphorylated by prior treatment with alkaline phosphatase. The 5' single-strand tail of the donor complex is complementary to the 12 nucleotides at the 5' end of the T7 transcript. Religation of the covalently held biotinylated DNA strand to the T7 transcript is observed as conversion of the 30-mer RNA to a product of 50 nucleotides.

Other Reference Publication (63):

Schmitt, et al., "Affinity purification of histidine-tagged proteins," Molecular Biology Reports 81:223-230 (1993).

ANSWER 1 OF 16 SCISEARCH COPYRIGHT 2004 THOMSON ISI on STN

ACCESSION NUMBER: 2004:195692 SCISEARCH

THE GENUINE ARTICLE: 774FM

TITLE: The DNA binding properties of the Escherichia

coli RecQ helicase

AUTHOR: Dou S X; Wang P Y; Xu H Q; Xi X G (Reprint)

CORPORATE SOURCE: Ecole Normale Super, CNRS, UMR 8113, Lab Biotechnol &

Pharmacol Genet Appl, 61 Ave President Wilson, F-94235 Cachan, France (Reprint); Ecole Normale Super, CNRS, UMR 8113, Lab Biotechnol & Pharmacol Genet Appl, F-94235 Cachan, France; Chinese Acad Sci, Inst Phys, Lab Soft

Matter Phys, Beijing 100080, Peoples R China

COUNTRY OF AUTHOR: France; Peoples R China

SOURCE: JOURNAL OF BIOLOGICAL CHEMISTRY, (20 FEB 2004) Vol. 279,

No. 8, pp. 6354-6363.

Publisher: AMER SOC BIOCHEMISTRY MOLECULAR BIOLOGY INC,

9650 ROCKVILLE PIKE, BETHESDA, MD 20814-3996 USA.

ISSN: 0021-9258. Article; Journal

DOCUMENT TYPE: LANGUAGE: English

REFERENCE COUNT: 52

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

The RecQ helicase family is highly conserved from bacteria to men and AB plays a conserved role in the preservation of genome integrity. Its deficiency in human cells leads to a marked genomic instability that is associated with premature aging and cancer. To determine the thermodynamic parameters for the interaction of Escherichia coli RecQ helicase with DNA, equilibrium binding studies have been performed

using the thermodynamic rigorous fluorescence titration technique. Steady-state fluorescence anisotropy measurements of fluorescein-labeled oligonucleotides revealed that RecQ helicase bound to DNA with an apparent binding stoichiometry of 1 protein monomer/10 nucleotides. This stoichiometry was not altered in the presence of AMPPNP (adenosine 5'-(beta,gamma-imido) triphosphate) or ADP. Analyses of RecQ helicase interactions with oligonucleotides of different lengths over a wide range of pH, NaCl, and nucleic acid concentrations indicate that the RecQ helicase has a single strong DNA binding site with an association constant at 25 degreesC of $K = 6.7 + / - 0.95 \times 10(6) M-1$ and a cooperativity parameter of omega = 25.5 +/- 1.2. Both single-stranded

DNA and double-stranded DNA bind competitively to the same site. The intrinsic affinities are salt-dependent, and the formation of DNA-helicase complex is accompanied by a net release of 3-4 ions. Allosteric effects of nucleotide cofactors on RecQ binding to DNA were observed only for single-stranded DNA in the

presence of 1.5 mm AMPPNP, whereas both AMPPNP and ADP had no detectable effect on double-stranded DNA binding over a large range of nucleotide cofactor concentrations.

ANSWER 2 OF 16 MEDLINE on STN DUPLICATE 1

ACCESSION NUMBER: 2003488948 MEDLINE DOCUMENT NUMBER: PubMed ID: 12909639

TITLE: RecQ helicase stimulates both DNA catenation and changes in

DNA topology by topoisomerase III.
Harmon Frank G; Brockman Joel P; Kowalczykowski Stephen C AUTHOR: - ----

Division of Biological Sciences, Section of Microbiology, Center for Genetics and Development, University of

California-Davis, 1 Shields Avenue, Davis, CA 95616, USA. Journal of biological chemistry, (2003 Oct 24) 278 (43)

42668-78.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

CORPORATE SOURCE:

SOURCE:

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

200401

ENTRY DATE:

Entered STN: 20031021

Last Updated on STN: 20040106 Entered Medline: 20040105

AB Together, RecQ helicase and topoisomerase III (Topo

III) of Escherichia coli comprise a potent DNA strand passage activity that can catenate covalently closed DNA (Harmon, F. G., DiGate, R. J., and Kowalczykowski, S. C. (1999) Mol. Cell 3, 611-620). Here we directly assessed the structure of the catenated DNA species formed by RecQ helicase and Topo III using atomic force microscopy. The images show complex catenated DNA species involving crossovers between multiple double-stranded DNA molecules that are consistent with full catenanes. E. coli single-stranded

DNA-binding protein significantly stimulated both the topoisomerase activity of Topo III alone and the DNA strand passage activity of RecQ helicase and Topo III. Titration data suggest that an intermediate of the RecQ helicase unwinding process, perhaps a RecQ helicase-DNA fork, is the target for Topo III action. Catenated DNA is the predominant product under conditions of molecular crowding; however, we also discovered that RecQ helicase and single-

stranded DNA-binding protein greatly stimulated the intramolecular strand passage ("supercoiling") activity of Topo III, as revealed by changes in the linking number of uncatenated DNA. Together our results demonstrate that RecQ helicase and Topo III function together to comprise a potent and concerted single-strand

DNA passage activity that can mediate both catenation-decatenation processes and changes in DNA topology.

L5 ANSWER 3 OF 16

MEDLINE on STN

DUPLICATE 2

ACCESSION NUMBER: 2001408241 DOCUMENT NUMBER: PubMed ID:

CORPORATE SOURCE:

2001408241 MEDLINE PubMed ID: 11459965

TITLE:

A yeast gene, MGS1, encoding a DNA-dependent AAA(+) ATPase

is required to maintain genome stability.

AUTHOR:

Hishida T; Iwasaki H; Ohno T; Morishita T; Shinagawa H Department of Molecular Microbiology, Research Institute

for Microbial Diseases, Osaka University, Yamadaoka 3-1,

Suita, Osaka 565-0871, Japan.

SOURCE:

Proceedings of the National Academy of Sciences of the United States of America, (2001 Jul 17) 98 (15) 8283-9.

Journal code: 7505876. ISSN: 0027-8424.

PUB. COUNTRY:

United States

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

200108

ENTRY DATE:

Entered STN: 20010903

Last Updated on STN: 20010903 Entered Medline: 20010830

AB Changes in DNA superhelicity during DNA replication are mediated primarily by the activities of DNA helicases and topoisomerases. If these activities are defective, the progression of the replication fork can be hindered or blocked, which can lead to double-strand breaks, elevated recombination in regions of repeated DNA, and genome instability.

Hereditary—diseases—like Werner's and Bloom's Syndromes are caused by defects in DNA helicases, and these diseases are associated with genome instability and carcinogenesis in humans. Here we report a Saccharomyces cerevisiae gene, MGS1 (Maintenance of Genome Stability 1), which encodes a protein belonging to the AAA(+) class of ATPases, and whose central region is similar to Escherichia coli RuvB, a Holliday junction branch migration motor protein. The Mgsl orthologues are highly conserved in prokaryotes and eukaryotes. The Mgsl protein possesses DNA-dependent ATPase and single-strand DNA annealing activities. An mgsl deletion mutant has an elevated rate of

mitotic recombination, which causes genome instability. The mgsl mutation is synergistic with a mutation in top3 (encoding topoisomerase III), and the double mutant exhibits severe growth defects and markedly increased genome instability. In contrast to the mgsl mutation, a mutation in the sgsl gene encoding a DNA helicase homologous to the Werner and Bloom helicases suppresses both the growth defect and the increased genome instability of the top3 mutant. Therefore, evolutionarily conserved Mgsl may play a role together with RecQ family helicases and DNA topoisomerases in maintaining proper DNA topology, which is essential for genome stability.

L5 ANSWER 4 OF 16 MEDLINE ON STN DUPLICATE 3

ACCESSION NUMBER: 2001372899 MEDLINE DOCUMENT NUMBER: PubMed ID: 11429611

TITLE: Crystal structure of a complex of a type IA DNA

topoisomerase with a single-stranded

DNA molecule.

AUTHOR: Changela A; DiGate R J; Mondragon A

CORPORATE SOURCE: Department of Biochemistry, Molecular Biology and Cell

Biology, Northwestern University, Evanston, Illinois 60208,

USA.

SOURCE: Nature, (2001 Jun 28) 411 (6841) 1077-81.

Journal code: 0410462. ISSN: 0028-0836.

PUB. COUNTRY: England: United Kingdom

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

OTHER SOURCE: PDB-117D ENTRY MONTH: 200107

ENTRY DATE: Entered STN: 20010723

Last Updated on STN: 20010723 Entered Medline: 20010719

AB A variety of cellular processes, including DNA replication, transcription, and chromosome condensation, require enzymes that can regulate the ensuing topological changes occurring in DNA. Such enzymes-DNA

topoisomerases-alter DNA topology by catalysing the cleavage of

single-stranded DNA (ssDNA) or double-stranded

DNA (dsDNA), the passage of DNA through the resulting break, and the rejoining of the broken phosphodiester backbone. DNA

topoisomerase III from Escherichia

coli belongs to the type IA family of DNA topoisomerases, which
transiently cleave ssDNA via formation of a covalent 5' phosphotyrosine
intermediate. Here we report the crystal structure, at 2.05 A resolution,
of an inactive mutant of E. coli DNA

topoisomerase III in a non-covalent complex with an 8-base ssDNA molecule. The enzyme undergoes a conformational change that allows the oligonucleotide to bind within a groove leading to the active site. We note that the ssDNA molecule adopts a conformation like that of B-DNA while bound to the enzyme. The position of the DNA within the realigned active site provides insight into the role of several highly conserved residues during catalysis. These findings confirm various aspects of the type IA topoisomerase mechanism while suggesting functional implications for other topoisomerases and proteins that perform DNA

rearrangements.

L5

ANSWER 5 OF 16 SCISEARCH COPYRIGHT 2004 THOMSON ISI on STN

ACCESSION NUMBER: 2000:688554 SCISEARCH

THE GENUINE ARTICLE: 351EL

TITLE: Interaction between yeast Sgsl helicase and DNA

topoisomerase III

AUTHOR: Bennett R J; NoirotGros M F; Wang J C (Reprint)

CORPORATE SOURCE: HARVARD UNIV, DEPT MOL & CELLULAR BIOL, 7 DIVIN AVE,

CAMBRIDGE, MA 02138 (Reprint); HARVARD UNIV, DEPT MOL &

CELLULAR BIOL, CAMBRIDGE, MA 02138

COUNTRY OF AUTHOR: USA

SOURCE: JOURNAL OF BIOLOGICAL CHEMISTRY, (1 SEP 2000) Vol. 275,

No. 35, pp. 26898-26905.

Publisher: AMER SOC BIOCHEMISTRY MOLECULAR BIOLOGY INC,

9650 ROCKVILLE PIKE, BETHESDA, MD 20814.

ISSN: 0021-9258.

DOCUMENT TYPE: Article; Journal

FILE SEGMENT: LIFE LANGUAGE: English

REFERENCE COUNT: 39

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

The Saccharomyces cerevisiae Sgs1 protein is a member of the RecQ family of DNA helicases that includes the human Bloom's syndrome and Werner's syndrome proteins. In this work, we report studies on the interaction between Sgs1 and DNA topoisomerase III in vitro and in vivo, Affinity chromatography experiments with various fragments of Sgs1, a 1447-amino acid polypeptide, suggested that its N-terminal one-fifth was sufficient for interaction with DNA topoisomerase III. Gel electrophoretic mobility shift assays also indicated that a fragment Sgs1(1-283), containing residues 1-283, inhibited the binding of DNA topoisomerase III to single-stranded DNA. A shorter protein fragment containing residues 1-107 also showed partial inhibition in these assays. Studies of a sgs1 top1 double mutant lacking both Sgs1 and DNA topoisomerase I showed that the slow growth phenotype of this double mutant is suppressed by expressing full-length Sqs1, but not Sqs1 without the N-terminal 107 amino acid residues. In sqsl top3 cells devoid of DNA topoisomerase III, however, expression of full-length Sqs1 or Sqs1 lacking the N-terminal 107 amino acid residues has the same effect of reducing the growth rate of the double mutant.

L5 ANSWER 6 OF 16 EMBASE COPYRIGHT 2004 ELSEVIER INC. ALL RIGHTS RESERVED.

on STN

ACCESSION NUMBER: 1999205900 EMBASE

TITLE: Purification and characterization of human DNA

These in vitro and in vivo data indicate that Sgs1 and DNA

topoisomerase III α .

topoisomerase III physically interact and that this

AUTHOR: Goulaouic H.; Roulon T.; Flamand O.; Grondard L.; Lavelle

F.; Riou J.-F.

interaction is physiologically significant.

CORPORATE SOURCE: H. Goulaouic, Rhone-Poulenc Rorer, Centre recherche

Vitry-Alfortville, 13 quai Jules Guesde, 94403 Vitry sur

Seine Cedex, France. helene.goulaouic@rp-rorer.fr

SOURCE: Nucleic Acids Research, (15 Jun 1999) 27/12 (2443-2450).

Refs: 31

ISSN: 0305-1048 CODEN: NARHAD

COUNTRY: United Kingdom DOCUMENT TYPE: Journal; Article

FILE SEGMENT: 022 Human Genetics

029 Clinical Biochemistry

LANGUAGE: English SUMMARY LANGUAGE: English

AB Human topoisomerase III α (hTopo III α), the

recently identified first member of the topoisomerase IA subfamily in humans, has a central domain which is highly homologous to the yeast topoisomerase III, but an overall organization closer to that of Escherichia coli DNA topoisomerase I. In order to determine the properties of hTopo III α , compared to those of other topoisomerase IA subfamily members, we purified this enzyme to near homogeneity, together with an active site-mutant Y337F. We show that hTopo III α is able to relax negatively supercoiled DNA in a distributive manner, leading to the total disappearance of the initial substrate and the appearance of intermediate topoisomers. This DNA relaxation activity

is magnesium-dependent, although a low concentration of MgCl2 is sufficient to obtain efficient catalysis. 32P-transfer experiments demonstrated that hTopo IIIα is able to cleave a single-stranded oligonucleotide and to bind covalently to the 5'-end of the cleaved DNA. Addition of 0.5 M NaCl reverses the reaction, leading to the religation of the oligonucleotide. Experiments utilizing several different single-stranded oligonucleotides permitted us to map several cleavage sites and to deduce a consensus sequence for DNA cleavage (CANNN↓), which is different from that for other members of the Topo IA subfamily.

L5 ANSWER 7 OF 16 MEDLINE on STN DUPLICATE 4

ACCESSION NUMBER: 2000045187 MEDLINE DOCUMENT NUMBER: PubMed ID: 10574789

TITLE: The structure of Escherichia coli DNA

topoisomerase III.

AUTHOR: Mondragon A; DiGate R

CORPORATE SOURCE: Department of Biochemistry, Molecular Biology and Cell

Biology, Northwestern University, Evanston, IL 60208-3500,

USA.. a-mondragon@nwu.edu

CONTRACT NUMBER: GM48445 (NIGMS)

GM51350 (NIGMS)

SOURCE: Structure with Folding & design, (1999 Nov 15) 7 (11)

1373-83.

Journal code: 100889329. ISSN: 0969-2126.

PUB. COUNTRY: ENGLAND: United Kingdom

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

OTHER SOURCE: PDB-1D6M ENTRY MONTH: 199912

ENTRY DATE: Entered STN: 20000113

Last Updated on STN: 20021030 Entered Medline: 19991214

Entered Medline: 19991214 AB BACKGROUND: DNA topoisomerases are enzymes that change the topology of DNA. Type IA topoisomerases transiently cleave one DNA strand in order to pass another strand or strands through the break. In this manner, they can relax negatively supercoiled DNA and catenate and decatenate DNA molecules. Structural information on Escherichia coli DNA topoisomerase III is important for understanding the mechanism of this type of enzyme and for studying the mechanistic differences among different members of the same subfamily. RESULTS: The structure of the intact and fully active E. coli DNA topoisomerase III has been solved to 3.0 A resolution. The structure shows the characteristic fold of the type IA topoisomerases that is formed by four domains, creating a toroidal protein. There is remarkable structural similarity to the 67 kDa N-terminal fragment of E. coli DNA topoisomerase I, although the relative arrangement of the four domains is significantly different. difference is the presence of a 17 amino acid insertion in topoisomerase III that protrudes from the side of the central hole and could be involved in the catenation and decatenation reactions. The active site is formed by highly conserved amino acids, but the structural information and existing biochemical and mutagenesis data are still insufficient to assign specific roles to most of them. -presence of a groove in one side of the protein is suggestive of a single-stranded DNA (ssDNA)-binding region. CONCLUSIONS: The structure of E. coli DNA topoisomerase III resembles the structure of E . coli DNA topoisomerase I except for the presence of a positively charged loop that may be involved in catenation and decatenation. A groove on the side of the protein leads to the active

site and is likely to be involved in DNA binding. The structure helps to

topoisomerases with greater confidence and expands the structural basis

establish the overall mechanism for the type IA subfamily of

for understanding these proteins.

ANSWER 8 OF 16 CAPLUS COPYRIGHT 2004 ACS on STN 1999:256931 CAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 131:112967 TITLE: Overexpression and purification of Escherichia coli DNA topoisomerase III AUTHOR (S): DiGate, Russell J. CORPORATE SOURCE: Department of Biomedical Chemistry, School of Pharmacy, University of Maryland at Baltimore, Baltimore, MD, USA SOURCE: Methods in Molecular Biology (Totowa, New Jersey) (1999), 94 (DNA Topoisomerase Protocols, Vol. 1), 153-162 CODEN: MMBIED; ISSN: 1064-3745 PUBLISHER: Humana Press Inc. DOCUMENT TYPE: Journal LANGUAGE: English A protocol for the purification of DNA topoisomerase III (topo III) of Escherichia coli is outlined. The procedure uses soybean trypsin inhibitor agarose, DE-52 cellulose batch, and single-stranded DNA cellulose chromatogs. and allows for purification of relatively large quantities of topo III in 3-4 day time period. THERE ARE 16 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 16 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT ANSWER 9 OF 16 SCISEARCH COPYRIGHT 2004 THOMSON ISI on STN ACCESSION NUMBER: 1998:311662 SCISEARCH THE GENUINE ARTICLE: ZH610 TITLE: Purification and characterization of the Sqs1 DNA helicase activity of Saccharomyces cerevisiae Bennett R J; Sharp J A; Wang J C (Reprint) AUTHOR: HARVARD UNIV, DEPT MOL & CELLULAR BIOL, 7 DIVIN AVE, CORPORATE SOURCE: CAMBRIDGE, MA 02138 (Reprint); HARVARD UNIV, DEPT MOL & CELLULAR BIOL, CAMBRIDGE, MA 02138 COUNTRY OF AUTHOR: USA JOURNAL OF BIOLOGICAL CHEMISTRY, (17 APR 1998) Vol. 273, SOURCE: No. 16, pp. 9644-9650. Publisher: AMER SOC BIOCHEMISTRY MOLECULAR BIOLOGY INC, 9650 ROCKVILLE PIKE, BETHESDA, MD 20814. ISSN: 0021-9258. DOCUMENT TYPE: Article; Journal FILE SEGMENT: LIFE LANGUAGE: English REFERENCE COUNT: 3.8 *ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS* The yeast Saccharomyces cerevisiae Sgs1 protein is a member of a family AB of DNA helicases that include the Escherichia coli RecQ protein and the products of human Bloom's syndrome and Werner's syndrome genes. To study the enzymatic characteristics of the protein, a recombinant Sgs1 fragment (amino acids 400-1268 of the 1447-amino acid full-length protein) was overexpressed in yeast and purified to near homogeneity. The purified protein exhibits an ATPase activity in the presence-of-single-or-double-stranded DNA. In the presence of ATP or dATP, unwinding of duplex DNA or a DNA-RNA heteroduplex by the recombinant Sgs1 fragment was readily observed. Similar to the E. coli RecQ helicase, displacement of the DNA strand occurs in the 3' to 5' direction with respect to the single-stranded DNA flanking the duplex. The efficiency of unwinding was found to

correlate inversely with the length of the duplex region and was enhanced

binding protein. In addition, the recombinant Sgsl fragment was found to bind more tightly to a forked DNA substrate than to either single-or

by the presence off. coli single-stranded DNA

double-stranded DNA.

ANSWER 10 OF 16 SCISEARCH COPYRIGHT 2004 THOMSON ISI on STN

ACCESSION NUMBER: 96:661945 SCISEARCH

THE GENUINE ARTICLE: VF614

TITLE: AN RNA TOPOISOMERASE

AUTHOR: WANG H; DIGATE R J; SEEMAN N C (Reprint)

CORPORATE SOURCE: NYU, DEPT CHEM, NEW YORK, NY, 10003 (Reprint); NYU, DEPT

CHEM, NEW YORK, NY, 10003; UNIV MARYLAND, MARYLAND

BIOTECHNOL INST, CTR MED BIOTECHNOL, BALTIMORE, MD, 21201; UNIV MARYLAND, MARYLAND BIOTECHNOL INST, DEPT PHARMACEUT

SCI, SCH PHARM, BALTIMORE, MD, 21201

COUNTRY OF AUTHOR:

SOURCE: PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE

UNITED STATES OF AMERICA, (03 SEP 1996) Vol. 93, No. 18,

pp. 9477-9482. ISSN: 0027-8424.

DOCUMENT TYPE: Article; Journal FILE SEGMENT: LIFE

LANGUAGE: ENGLISH REFERENCE COUNT:

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

A synthetic strand of RNA has been designed so that it can adopt two different topological states (a circle and a trefoil knot) when ligated into a cyclic molecule. The RNA knot and circle have been characterized by their behavior in gel electrophoresis and sedimentation experiments, This system allows one to assay for the existence of an RNA topoisomerase, because the two RNA molecules can be interconverted only by a strand passage event. We find that the interconversion of these two species can be catalyzed by Escherichia coli DNA

topoisomerase III, indicating that this enzyme can act as an RNA topoisomerase. The conversion of circles to knots is accompanied by a small amount of RNA catenane generation. These findings suggest that strand passage must be considered a potential component of the folding and modification of RNA structures.

ANSWER 11 OF 16 DUPLICATE 5 MEDLINE on STN

ACCESSION NUMBER: 96224127 MEDLINE DOCUMENT NUMBER: PubMed ID: 8621552 DOCUMENT NUMBER:

The role of the carboxyl-terminal amino acid residues in TITLE:

Escherichia coli DNA

topoisomerase III-mediated catalysis.

Zhang H L; Malpure S; Li Z; Hiasa H; DiGate R J AUTHOR:

Molecular and Cell Biology Program, University of Maryland, Baltimore, Maryland 21201, USA. CORPORATE SOURCE:

CONTRACT NUMBER: GM48445 (NIGMS)

SOURCE: Journal of biological chemistry, (1996 Apr 12) 271 (15)

9039-45.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199606

ENTRY_DATE: ____Entered STN: 19960627

Last Updated on STN: 19960627 Entered Medline: 19960620

AB The role that the carboxyl-terminal amino acids of Escherichia coli DNA topoisomerase I (Topo I) and III (Topo III) play in catalysis was examined by comparing the properties of Topo III with those of a truncated enzyme lacking the generalized DNA binding domain of Topo III, Topo I, and a hybrid topoisomerase polypeptide containing the amino-terminal 605 amino acids of Topo III and the putative generalized DNA binding domain of Topo I. The deletion of the carboxyl-terminal 49

amino acids of Topo III decreases the affinity of the enzyme for its substrate, **single-stranded DNA**, by

approximately 2 orders of magnitude and reduces Topo III-catalyzed relaxation of supercoiled DNA and Topo III-catalyzed resolution of DNA replication intermediates to a similar extent. Fusion of the carboxyl-terminal 312 amino acid residues of Topo I onto the truncated molecule stimulates topoisomerase-catalyzed relaxation 15-20-fold, to a level comparable with that of full-length Topo III. However, topoisomerase-catalyzed resolution of DNA replication intermediates was only stimulated 2-3-fold. Therefore, the carboxyl-terminal amino acids of these topoisomerases constitute a distinct and separable domain, and this domain is intimately involved in determining the catalytic properties of these polypeptides.

L5 ANSWER 12 OF 16 MEDLINE on STN DUPLICATE 6

ACCESSION NUMBER: 96007520 MEDLINE
DOCUMENT NUMBER: PubMed ID: 7559540
TITLE: Escherichia coli DNA

topoisomerase III is a site-specific DNA

binding protein that binds asymmetrically to its cleavage

site.

AUTHOR: Zhang H L; Malpure S; DiGate R J

CORPORATE SOURCE: Molecular and Cell Biology Program, University of Maryland

at Baltimore 21201, USA.

CONTRACT NUMBER: GM48445-02 (NIGMS)

SOURCE: Journal of biological chemistry, (1995 Oct 6) 270 (40)

23700-5.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY: United States

DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals

ENTRY MONTH: 199511

ENTRY DATE: Entered STN: 19951227

Last Updated on STN: 19951227 Entered Medline: 19951114

AB The binding of DNA topoisomerase III (Topo III) to a

single-stranded DNA substrate containing a strong cleavage site has been examined. The minimal substrate requirement for Topo III-catalyzed cleavage has been determined to consist of 7 bases; 6 bases 5' to the cleavage site and only 1 base 3' to the site. Nuclease P1 protection experiments indicate that the enzyme also binds to its substrate asymmetrically, protecting approximately 12 bases 5' to the cleavage site and only 2 bases 3' to the cleavage site. A catalytically inactive mutant of Topo III shows the same protection pattern as the active polypeptide, indicating that Topo III is a site-specific binding protein as well as a topoisomerase. Consistent with this view, an oligonucleotide containing a cleavage site is a more effective inhibitor and is bound more efficiently by Topo III than an oligonucleotide without a cleavage site.

L5 ANSWER 13 OF 16 MEDLINE ON STN DUPLICATE 7

ACCESSION NUMBER: 94179321 MEDLINE DOCUMENT NUMBER: PubMed ID: 7510701

TITLE: --- -- The carboxyl-terminal residues of Escherichia

coli DNA topoisomerase III are involved in substrate binding.

COMMENT: Erratum in: J Biol Chemical 1995 Sep 1;270(35):20870. PubMed

ID: 7657674

AUTHOR: Zhang H L; DiGate R J

CORPORATE SOURCE: Molecular and Cell Biology Program, University of Maryland

at Baltimore 21201.

CONTRACT NUMBER: GM48445 (NIGMS)

RR05770-14 (NCRR)

SOURCE: Journal of biological chemistry, (1994 Mar 25) 269 (12)

9052-9.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY:

United States

Journal; Article; (JOURNAL ARTICLE) DOCUMENT TYPE:

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

199404

ENTRY DATE:

Entered STN: 19940428

Last Updated on STN: 19960315

Entered Medline: 19940421

AB The nucleic acid-binding domain of Escherichia coli DNA topoisomerase III (Topo III) has been identified

using a selection procedure designed to isolate inactive Topo III polypeptides. Deletion of this binding domain, contained in the carboxyl terminus of Topo III, results in a drastic reduction in the ability of the enzyme to bind to single-stranded DNA and

RNA substrates. Successive truncation of the enzyme within this region results in the gradual loss of nucleic acid binding activity and in a gradual change in the mechanism of Topo III-catalyzed relaxation of negatively supercoiled DNA. The reduction of nucleic acid binding activity of the truncated polypeptides does not result in a loss of cleavage site specificity for the enzyme, suggesting that other amino acids are involved in the positioning of the nucleic acid within the nicking/closing site of the topoisomerase.

ANSWER 14 OF 16 MEDLINE on STN **DUPLICATE 8**

ACCESSION NUMBER: 92381032

MEDLINE

DOCUMENT NUMBER:

PubMed ID: 1324925

TITLE:

Identification of the yeast TOP3 gene product as

a single strand-specific DNA topoisomerase.

AUTHOR:

Kim R A; Wang J C

CORPORATE SOURCE:

Department of Biochemistry and Molecular Biology, Harvard

University, Cambridge, Massachusetts 02138-2092.

CONTRACT NUMBER:

GM 24544 (NIGMS)

SOURCE:

Journal of biological chemistry, (1992 Aug 25) 267 (24)

17178-85.

Journal code: 2985121R. ISSN: 0021-9258.

PUB. COUNTRY:

United States

CA47958 (NCI)

DOCUMENT TYPE:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE:

English

FILE SEGMENT:

Priority Journals

ENTRY MONTH:

199209

ENTRY DATE:

Entered STN: 19921018

Last Updated on STN: 19921018 Entered Medline: 19920925

The TOP3 gene of the yeast Saccharomyces cerevisiae was

postulated to encode a DNA topoisomerase, based on its sequence homology

to Escherichia coli DNA topoisomerase I and the

suppression of the poor growth phenotype of top3 mutants by the

expression of the E. coli enzyme (Wallis, J.W.,

Chrebet, G., Brodsky, G., Golfe, M., and Rothstein, R. (1989) Cell 58,

409-419). We have purified the yeast TOP3 gene product to near

homogeneity as a 74-kDA-protein from yeast cells lacking DNA topoisomerase

I and overexpressing a plasmid-borne TOP3 gene linked to a

phosphate-regulated yeast PHO5 gene promoter. The purified protein possesses a distinct DNA topoisomerase activity: similar to E.

coli DNA topoisomerases I and III, it partially relaxes negatively

but not positively supercoiled DNA. Several experiments, including the use of a negatively supercoiled heteroduplex DNA containing a

29-nucleotide single-stranded loop, indicate that the activity has a strong preference for single-stranded DNA.

A protein-DNA covalent complex in which the 74-kDa protein is linked to a

5' DNA phosphoryl group has been identified, and the nucleotide sequences of 30 sites of DNA-protein covalent complex formation have been determined. These sequences differ from those recognized by E. coli DNA topoisomerase I but resemble those recognized by

E. coli DNA topoisomerase III.

Based on these results, the yeast TOP3 gene product can formally be termed S. cerevisiae DNA topoisomerase III.

Analysis of supercoiling of intracellular yeast plasmids in various DNA topoisomerase mutants indicates that yeast DNA topoisomerase III has at most a weak activity in relaxing negatively supercoiled double-stranded DNA in vivo, in accordance with the characteristics of the

purified enzyme.

ANSWER 15 OF 16 SCISEARCH COPYRIGHT 2004 THOMSON ISI on STN

ACCESSION NUMBER: 91:380018 SCISEARCH

THE GENUINE ARTICLE: FV180

TITLE:

DNA TOPOISOMERASE-III FROM EXTREMELY

THERMOPHILIC ARCHAEBACTERIA - ATP-INDEPENDENT TYPE-I TOPOISOMERASE FROM DESULFUROCOCCUS-AMYLOLYTICUS DRIVES

EXTENSIVE UNWINDING OF CLOSED CIRCULAR DNA AT

HIGH-TEMPERATURE

SLESAREV A I (Reprint); ZAITZEV D A; KOPYLOV V M; STETTER AUTHOR:

K O; KOZYAVKIN S A

UNIV REGENSBURG, LEHRSTUHL MIKROBIOL, UNIV STR 31, W-8400 CORPORATE SOURCE:

REGENSBURG, GERMANY (Reprint); ACAD SCI USSR, INST MOLEC GENET, MOSCOW 123182, USSR; RE KAVETSKY ONCOL PROBLEMS

INST, KIEV 252127, UKRAINE, USSR

GERMANY; USSR COUNTRY OF AUTHOR:

SOURCE:

JOURNAL OF BIOLOGICAL CHEMISTRY, (1991) Vol. 266, No. 19,

pp. 12321-12328.

DOCUMENT TYPE:

Article; Journal

FILE SEGMENT: LANGUAGE:

LIFE ENGLISH

REFERENCE COUNT: 73

discussed.

ABSTRACT IS AVAILABLE IN THE ALL AND IALL FORMATS

A second type I topoisomerase was purified from the extremely AR thermophilic archaebacterium Desulfurococcus amylolyticus. In contrast to the previously described reverse gyrase from this organism, the novel enzyme designated as Dam topoisomerase III is an ATP-independent relaxing topoisomerase. It is a monomer with M(r) 108,000, as determined by electrophoresis under denaturing conditions and by size exclusion chromatography. Dam topoisomerase III, like other bacterial type I topoisomerases, absolutely requires Mg2+ for activity and is specific for single-stranded DNA. At 60-80-degrees-C, it relaxes negatively but not positively supercoiled DNA and is inhibited by single-stranded M13 DNA. At 95-degrees-C, the enzyme unwinds both positively and negatively supercoiled substrates and produces extensively unwound form I* and I**

DNA. The peculiarities of DNA topoisomerization at high temperatures are

ANSWER 16 OF 16 MEDLINE on STN DUPLICATE 9

ACCESSION NUMBER: 84203564 MEDLINE DOCUMENT NUMBER: PubMed ID: 6326814 DOCUMENT NUMBER: TITLE: ____Escherichia coli DNA

topoisomerase III: purification and

characterization of a new type I enzyme. AUTHOR: Srivenugopal K S; Lockshon D; Morris D R Biochemistry, (1984 Apr 24) 23 (9) 1899-906. SOURCE:

Journal code: 0370623. ISSN: 0006-2960.

United States

PUB. COUNTRY:
DOCUMENT TYPE: Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

FILE SEGMENT: Priority Journals ENTRY MONTH:

198407

topoisomerase activity.

ENTRY DATE:

Entered STN: 19900319

Last Updated on STN: 19970203

Entered Medline: 19840711

A new topoisomerase capable of relaxing negatively supercoiled DNA in Escherichia coli has been identified during chromatography on novobiocin-Sepharose. A simple and reproducible purification procedure is described to obtain this enzyme, called topoisomerase III (topo III), in a homogeneous form. The protein is a single polypeptide with a molecular weight of 74 000 +/-2000 and is a type I topoisomerase, changing the linking number of DNA circles in steps of one. It is present in deletion strains lacking the topA gene and further differs from the well-studied topoisomerase I (omega protein; Eco topo I) in (1) its requirement for K+ in addition to Mg2+ to exhibit optimal activity and (2) its affinity to novobiocin-Sepharose. Positively supercoiled DNA is not relaxed during exposure to the enzyme. Topo III has no ATPase activity, and ATP does not show any discernible effect on the reduction of superhelical turns. The purified topoisomerase has no supercoiling activity and is unaffected by high concentrations of oxolinic acid and novobiocin in the relaxing reaction. Singlestranded DNA and spermidine strongly inhibit the